PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

INTERNATIONAL APPLICATION PUBLIS	MED	JNDER THE PATENT COOPERATION TREATT (FCT)
(51) International Patent Classification ³ :		(11) International Publication Number: WO 82/ 02719
C08L 33/12; C09D 3/81	A1	(43) International Publication Date: 19 August 1982 (19.08.82)
(21) International Application Number: PCT/AU (22) International Filing Date: 4 February 1982 (Street, P.O. Box 4311, Melbourne, vic. 3001 (AU).
(31) Priority Application Number: (32) Priority Date: 13 February 1981 (PE 75 (13.02.8	pean patent), DK, FR (European patent), GB (Euro-
(33) Priority Country:	A	.U
(71) Applicant: DULUX AUSTRALIA LTD. [AU Collins Street, Melbourne, Vic. 3000 (AU).	ī/AU];	Published With international search report.
(72) Inventors: HALL, Derrard, Michael; Snowdo Glen Waverley, Vic. 3150 (AU). KEMP, Roy 41 Glenelg Road, Fernglen, Port Elizabeth, Cvince (ZA). SUCH, Christopher, Henry; 86 Avenue, Mount Eliza, Vic. 3930 (AU).	, Arthu Cape Pr	r ; o-

(54) Title: ACRYLIC COATING COMPOSITIONS

(57) Abstract

An acrylic coating composition suitable for use as an automotive refinish lacquer comprises a mixture of (a) a solution in volatile organic solvent of an acrylic copolymer of 75 - 90 % methyl methacrylate, 1,5 - 6,0 % dimethylaminoethyl methacrylate and 8 - 23,5 % of butyl acrylate, butyl methacrylate or lauryl methacrylate, the copolymer having a Tg of 80 - 95 °C; (b) a solution in volatile organic solvent of cellulose acetate butyrate, the cellulose acetate butyrate having a viscosity of 0,5 - 5 sec and a hydroxyl value of 2 maximum; and (c) a monomeric ester plasticiser such that the non-volatile constituents of (a), (b) and (c) comprise respectively 43,5 - 64,0 %, 15,0 - 25,5 % and 2,10 - 31,5 % of the total weight of non-volatile material of (a) plus (b) plus (c).

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	KP	Democratic People's Republic of Korea
AU	Australia	LI	Liechtenstein
BE	Belgium	LK	Sri Lanka
BR	Brazil	LU	Luxembourg
CF	Central African Republic	МC	Мопасо
CG	Congo	MG	Madagascar
CH	Switzerland	MW	Malawi
CM	Cameroon	NL	Netherlands
DE	Germany, Federal Republic of	NO	Norway ·
DK	Denmark	RO	Romania
FI	Finland	SE	Sweden
FR	France	SN	Senegal
GA	Gabon	SU	Soviet Union
GB	United Kingdom	TD	Chad
HU	Hungary	TG	Togo
JP	Japan	US	United States of America

ACRYLIC COATING COMPOSITIONS

This invention relates to thermoplastic coating compositions and in particular to such compositions in which the principal film-forming ingredients are three-component blends of acrylic polymers, cellulose acetate butyrate and monomeric plasticiser.

According to the present invention we now provide a liquid coating composition comprising a mixture of

- (a) a solution in volatile organic liquid of a copolymer of from 75-90% methyl methacrylate,
- 1.5 4.0% dimethyl aminoethyl methacrylate and 8 23.5% of monomer selected from butyl methacrylate, butyl acrylate and lauryl methacrylate, which copolymer shall have a glass transition temperature of 80° 95°C;
- 15 (b) a solution in volatile organic liquid of cellulose acetate butyrate which has a viscosity of 0.5 5.0 sec when measured according to Test Method D1343 of the American Society for Testing and Materials, and a hydroxyl value of 2 maximum; and
- 20 (c) a monomeric ester plasticiser compatible with the selected mixture of the said components (a) plus (b)
 - such that the non-volatile constituents of (a), (b) and (c) comprise respectively 43.5 64.0%, 15.0 25.5%
- and 21.0 31.5% of the total weight of non-volatile material in (a) plus (b) plus (c).

We further provide a process of preparing a liquid coating composition by blending

(d) a solution in volatile liquid solvent of a copolymer of from 75-90% methyl methacrylate, 1.5 - 4.0% dimethyl aminoethyl methacrylate and 8 - 23.5% of monomer selected from butyl methacrylate, butyl acrylate and lauryl methacrylate, which copolymer shall have a glass transition temperature of 80° - 95°C;



- (e) a solution in volatile liquid solvent of cellulose acetate butyrate which has a viscosity of 0.5 - 5.0 sec when measured according to Test Method D1343 of the American Society for Testing and Materials, and a hydroxyl value of 2 maximum; and
- (f) a monomeric ester plasticiser compatible with the selected mixture of the said components (d) plus (e)
- such that the non-volatile constituents of (d), (e) and (f) comprise respectively 43.5 64.0%, 15.0 25.5% and 21.0 31.5% of the total weight of non-volatile material in (d) plus (e) plus (f).
- The compositions of this invention provide the

 15 principal film-forming components of liquid coating
 compositions, which are particularly suitable for use
 as automotive lacquers. Paint films deposited therefrom exhibit particularly useful combinations of
 hardness, flexibility, gloss and in particular,
 20 adhesion to other paint substrates.

The composition range of components which will confer these desirable properties is unusually narrow and to confer the optimum combination of hardness and adhesion for some uses we have found that it must be limited even further to the following preferred limits.

The preferred liquid coating composition comprises a mixture of

(a) a solution in volatile organic liquid of a copolymer of from 75-80% methyl methacrylate, 2.5 - 3.5% dimethyl aminoethyl methacrylate and 1.5 - 2.0% butyl methacrylate, which shall have a glass transition temperature of 80° - 95°C.

> OMPI WIFO

30

25

10

25

- (b) a solution in volatile organic liquid of cellulose acetate butyrate which has a viscosity of 0.5 - 5.0 sec when measured according to Test Method D1343 of the American Society for Testing and Materials, and which has a hydroxyl value of 2 maximum; and
- (c) a compatible monomeric ester plasticisers) selected from butyl benzyl phthalate and dipropylene glycol dibenzoate.

such that the non-volatile constituents of (a), (b) and (c) comprise respectively 43.5 - 64.0%, 15.0 - 25.5% and 21.0 - 31.5% of the total weight of non-volatile material in (a) plus (b) plus (c)

The preparation of component (a) is a conventional copolymerisation process which may be performed using standard polymerisation techniques. The glass transition temperature may be measured by using, for example a Differential Thermal Mechanical Aralyser, but for the purpose of determining compliance with our invention, it is satisfactory to calculate the value from the known composition using published Tg values for homopolymers of the individual components.

As is usual with liquid coating composition, these will normally contain from 60-80% by weight of volatile solvent.

Pigments may also be incorporated in the coatings in conventional manner.

The invention is illustrated by the following examples, in which all components are expressed by weight:



20

25

30

35

EXAMPLE 1

Preparation of a coating composition according to the invention and comparative testing against a commercial lacquer not according to this invention.

An acrylic copolymer solution of the following composition was prepared by conventional addition polymerisation techniques:

monomer composition

		-	
		methyl methacrylate	83 parts
10		n-butyl methacrylate	15 "
		dimethyl aminoethyl	
		methacrylate	2"
	solids		40% by weight
	solvent	composition	-
15		methyl ethyl ketone	30 parts
		toluene	70 "

132.5 parts of this solution was blended with 88 parts of a 25% (by weight) solution of cellulose acetate butyrate (0.5 second grade) in methyl ethyl ketone and toluene (30 : 70 by weight) and 25.0 parts of n-butyl benzyl phthalate, to give a solution of film-forming polymer wherein the non-volatile constituents of the copolymer solution, the cellulose acetate butyrate solution and the plasticiser comprised 53%, 22% and 25% of the weight of the total non-volatile content.

The solution so-prepared was pigmented to a pigment/binder ratio of 60 with titanium dioxide pigment to produce a white lacquer.

This lacquer was tested according to the methods outlined below. A commercial acrylic refinish lacquer having the same overall general composition but in which the acrylic polymer of this example was replaced with an equal weight, based on solids content, of a methyl methacrylate homopolymer was also tested according to these methods.



10

15

20

25

30

٢

Adhesion test_

The lacquer was thinned with the abovementioned solvent to a viscosity of 15-17 sec. in a Ford 4 viscosity cup and sprayed on to steel test panels coated with a baked, aged and lightly sanded film of commercial thermoset acrylic enamel. The film was dried in air for 24 hours and then subjected to the adhesion test of the American Society for Testing and Materials, test method D3359.

Heat distortion test

The lacquer was applied to a primed steel panel and the panel was spotted along its length with a 1% solution of egg albumen. The spots were allowed to dry overnight and the panel was then placed for 6 minutes on a thermal copper bar with a temperature gradient of from 35°C to 65°C between its ends. The panel was washed and inspected for distortion resistance, the temperature to which each albumen spot was heated being easily determined by its position on the bar. A coating composition is considered to have acceptable distortion resistance if only those spots corresponding to temperatures above about 40°C show signs of distortion.

Recoat craze test

The lacquer was applied to a primed steel panel, allowed to air dry and aged in an oven at 65°C for 16 hours. The panel was then cooled to 25°C and a 175 µm film of the lacquer (thinned to spraying viscosity) was drawn down. The film was allowed to dry and examined for crazing (a network of fine interconnecting cracks on the surface of the dry film). The panel was returned to the oven for seven days after which it was again withdrawn, cooled to 25°C and the abovementioned procedure repeated. The panel and



15

20

25

30

the thinned lacquer used for the draw-downs were then cooled to 7°C and a third film drawn down, allowed to dry and examined for crazing.

Both lacquers performed satisfactorily in the heat distortion and recoat craze test but whereas the lacquer prepared according to the invention exhibited almost no loss of adhesion in the adhesion test, the commercial lacquer exhibited very poor adhesion.

10 EXAMPLE 2

Use of a number of different acrylic polymers.

A number of coating compositions was prepared according to Example 1 but in each case the copolymer in that example was replaced by one of the following copolymers;

- (a) methyl methacrylate (87%), n-butylmethacrylate (10%), dimethylaminoethylmethacrylate (DMAEMA) (3%).
- (b) methyl methacrylate (92%), laurylmethacrylate (6%), DMAEMA (2%);
- (c) methyl methacrylate (92%), butylacrylate (6%), DMAEMA (2%);
- (d) methyl methacrylate (90%), ethylacrylate (8%), DMAEMA (2%);
- (e) methyl methacrylate (95%), n-butyl methacrylate (3%), DMAEMA (2%)
- where (a) (b) (c) are according to the invention, (d) includes a monomer which is not according to the invention and (e) has proportions of comonomers which lie outside the permitted limits.



25

30

- 7 -

The coating compositions thus prepared were subjected to the testing procedures of Example 1. The compositions (a), (b) and (c) gave good results in all tests whereas the composition (d) performed poorly in the adhesion and heat distortion tests and the composition (e) in the adhesion and recoat craze test.

EXAMPLE 3

Effect of changing the proportions of the 10 components of the coating composition.

Example 1::was repeated but the proportions of the components were altered such that the ratio of copolymer/cellulose acetate butyrate/placticiser was 69/6/25 instead of the 53/22/25 ratio of that example.

When subjected to the testing procedures outlined in Example 1 the resultant coating composition exhibited unacceptable recoat crazing.

EXAMPLE 4

Use of various types of cellulose acetate 20 butyrate (CAB).

Example 1 was repeated twice, in each case replacing the CAB of that example by an identical quantity of

- (a) a CAB whose hydroxyl value was 3; and
- (b) a CAB whose viscosity was 0.2 sec.

When tested by the procedures outlined in Example 1, the composition containing (a) exhibited poor adhesion and that containing (b) poor recoat crazing resistance. Prior to testing, it was also noticeable that each composition had poor gloss when compared to the composition of Example 1.



EXAMPLE 5

Use of various plasticisers.

Example 1 was repeated, replacing the butyl benzyl phthalate plasticiser of that example by an identical quantity of

- (a) dipropylene glycol dibenzoate
- (b) "Santiciser" 141 (trade mark) ex Monsanto Co. - a commercially available monomeric plasticiser
- 10 (c) 34% coconut glycerol alkyd resin
 - (d) saturated neo-pentylglycol-adipic acid polyester.

The compositions containing (a) and (b) performed satisfactorily when tested according to the procedures of Example 1 but those containing (c) and (d) exhibited poor adhesion.

Claims

- 1. A liquid coating composition comprising a mixture of
 - (a) a solution in volatile organic liquid of a copolymer of from 75 90% methyl methacrylate, 1.5 4.0% dimethyl aminoethyl methacrylate and 8 23.5% of monomer selected from butyl methacrylate, butyl acrylate and lauryl methacrylate, which copolymer shall have a glass transition temperature of 80° 95°C;
 - (b) a solution in volatile organic liquid of cellulose acetate butyrate which has a viscosity of 0.5 5.0 sec when measured according to Test Method D1343 of the American Society for Testing and Materials, and a hydroxyl value of 2 maximum; and
 - (c) a monomeric ester plasticiser compatible with the selected mixture of the said components (a) plus (b)

such that the non-volatile constituents of (a), (b) and (c) comprise respectively 43.5 - 64.0%, 15.0 - 24.5% and 21.0 - 31.5% of the total weight of non-volatile material of (a) plus (b) plus (c)

- 2. A liquid coating composition according to claim 1 wherein (a) the copolymer comprises 75 80% methyl methacrylate, 2.5 3.5% dimethylaminoethyl methacrylate and 1.5 2.5% butyl methacrylate; and (b) the monomeric ester plasticiser is selected from butyl benzyl phthalate and dipropylene glycol dibenzoate.
- 3. A process of preparing a liquid coating composition by blending
 - (d) a solution in volatile liquid solvent of a copolymer of from 75 90% methyl methacrylate.
 1.5 4.0% dimethyl aminoethyl methacrylate and 8 23.5% of monomer selected from butyl methacrylate, butyl acrylate and lauryl methacrylate, which copolymer shall have a glass transition temperature of 80° 95°C;



- (e) a solution in volatile liquid solvent of cellulose acetate butyrate which has a viscosity of 0.5 - 5.0 sec when measured according to Test Method D1343 of the American Society for Testing and Materials, and a hydroxyl value of 2 maximum; and
- (f) a monomeric ester plasticiser compatible with the selected mixture of the said components (d) plus (e)

such that the non-volatile constituents of (d), (e) and (f) comprise respectively 43.5 - 64.0%, 15.0 - 25.5% and 21.0 - 31.5% of the total weight of non-volatile material of (d) plus (e) plus (f)

A process according to claim 3 wherein

- (d) the copolymer comprises 75 80% methyl methacrylate, 2.5 3.5% dimethylaminoethyl methacrylate and 1.5 2.5% butyl methacrylate; and
- (e) the monomeric ester plasticiser is selected from butyl benzyl phthalate and dipropylene glycol dibenzoate.



INTERNATIONAL SEARCH REPORT PCT/AU82/00008 International Application No 1. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) 1 According to International Patent Classification (IPC) or to both National Classification and IPC Int. Cl. 3 COSL 33/12 CO9D 3/81 II. FIELDS SEARCHED Minimum Documentation Searched Classification Symbols Classification System COSL 33/12 · · · IPC 106;169, 260;17R, 525;186 US C1 Documentation Searched other than Minimum Documentation to the Extent that such Occuments are included in the Fields Searched 5 AU: IPC as above; Australian Classification 47.7_481 III. DOCUMENTS CONSIDERED TO BE RELEVANT 14 Relevant to Claim No. 15 Citation of Document, 16 with Indication, where appropriate, of the relevant passages 17 Category ⁴ AU, B, 62152/69 (432012) E.I. DUPONT DE NEMOURS AND 1-4 Α COMPANY) 22 April 1971 (22.04.71) US,A, 3637546 (PARKER) 25 January 1972 (25.01.72) 1-4 Α US,A, 3862062 (HARPER) 21 January 1975 (21.01.75) 1-4 A US,A, 3862063 (PETTIT) 21 January 1975 (21.01.75) 1-4 Α US,A, 4279789 (LUEDDECKE et al) 21 July 1981 1-4 A (21.07.81)EP,A, 29594 (E.I. DUPONT DE NEMOURS AND COMPANY 1-4 A 3 June 1981 (03.06.81). "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the Special categories of cited documents: 15 "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is compined with one or more other such documents, such combination being obvious to a person skilled in the control of th "O" document referring to an oral disclosure, use, exhibition or ments, sur in the art. other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family IV. CERTIFICATION Date of Mailing of this International Search Report 3 Date of the Actual Completion of the International Search 3

se April

D.B. CUPITT

/C 6 C 4 SZ)_
Signature of Authorized Officer 10

1932

Australian Patent Office
Form PCT/ISA/210 (second sheet) (October 1981)

International Searching Authority 1

31 March 1982, (31.03.82)